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Applicant: Duncan Roger HARPER, et al.

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Serial No.: 10/091,284

Examiner: Unknown

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: March 5, 2002

Title

: AEROSOL COMPOSITIONS TO WHICH AN ELECTROSTATIC CHARGE

CAN BE IMPARTED UPON ACTUATION

Commissioner for Patents Washington, D.C. 20231

CLAIM OF PRIORITY UNDER 35 USC §119

Sir:

Duncan Roger HARPER, Neale HARRISON, John Douglas MORGAN, John Howard CLINT and Mario ABELA, applicants in this application, hereby claim the priority date of British Application No. 9921037.9 filed on 7 September 1999. A certified copy of said British Application is here enclosed.

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Respectfully submitted,

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Dated 5 June 2002

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Title of the invention

COMPOSITIONS

Name of your agent (if you have one)

Patents ADP number (if you know it)

country/state of its incorporation

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Elizabeth A Dickson

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If you are declaring priority from one or more earlier patent applications, give the country and the date of filing of the or of each of these earlier applications and (if you know it) the or each application number

Country

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Is a statement of inventorship and of right to grant of a patent required in support of this request? (Answer 'Yes' if:

any applicant named in part 3 is not an inventor, or

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I/We request the grant of a patent on the basis of this application.

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Date 7 September 1999

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 Name and daytime telephone number of Person to contact in the United Kingdom

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COMPOSITIONS BY CORP. STATE OF STATE OF

enged th The Control of Control The present invention relates to aerosol compositions and, in particular, compositions in which the droplets are imparted with an electrostatic charge on spraying from an aerosol spray device and in which the electrostatic charge on the droplets is maximised through the inclusion in the compositions of certain selected components.

Aerosol spray devices are a convenient form in which a variety of useful products, such as insecticides, air fresheners, antiperspirants, hair. sprays, horticultural products, waxes and polishes, oven cleaners, starches and fabric finishes, shoe and leather care products, glass cleaners and various and other household, institutional, professional or industrial products, can be dispensed...

The utility of aerosol-spray devices resides in ... the ability to readily deliver the composition contained within the device in the form of fine droplets to the target area, for example the spraying of an insecticide onto target insects.

In WO 97/28883 there is described a method of precipitating airborne particles from air in a domestic environment containing such particles in which the air to be treated is sprayed with liquid droplets from an aerosol spray device with a unipolar charge being imparted to the droplets during the spraying of the liquid droplets by the aerosol spray - ** device, the unipolar charge being atwa level such that 🕔 . the droplets have a charge to mass ratio of at least " " $+/- 1 \times 10^{-4} \text{ C/kg}.$

In WO 99/01227 there is described a method of killing flying insects by spraying into the air in which the insects are flying liquid droplets of an

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insecticidal composition, a unipolar charge being imparted to the liquid droplets by double layer charging and charge separation during spraying, the unipolar charge being at a level such that the said liquid droplets have a charge to mass ratio of at least +/- 1 x 10⁻⁴ C/kg. An apparatus for imparting the unipolar charge of this magnitude to a liquid composition is also described.

We have now found that by careful selection of the components which are to be contained within a liquid composition for application by aerosol spraying, it is possible to charge the liquid droplets during the spraying operation without requiring any special features of the construction of the aerosol spraying head.

Accordingly, in one aspect the present invention provides an electrically neutral composition in the form of a water-in-oil or an oil-in-water emulsion, in which droplets of the emulsion on discharge from an aerosol spray device are imparted with a unipolar electrostatic charge, which composition comprises:

- (a) at least one propellant in an amount of from 2 to 80% w/w;
- (c) optionally one or more solvents within the oil phase in an amount of up to 40% $\rm w/v_{\odot}$
- (d) at least one polar or ionic or aromatic or conjugated compound which is attracted to the interface between the disperse phase and the continuous phase of the emulsion in an amount of from 0.01 to 18% w/w based on the non-ionic surfactant, but which is such that the theoretical conductivity of the emulsion is less than the bulk conductivity of the emulsion; and

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In a second aspect the present invention provides a method of enhancing the unipolar charge which is imparted to droplets of an emulsion on discharge from an aerosol spray device in which the droplets are formed from an oil-in-water or a water-in-oil emulsion composition which comprises:

- (a) at least one propellant in an amount of from 2 to 80% w/w; and the propellant in the amount of from
- (b) at least one non-ionic surfactant in an amount of from 0.01 to 10% w/w;
- oil phase in an amount of up to 40% w/w;
- (d) at least one polar or ionic or aromatic or conjugated compound which is attracted to the interface between the disperse phase and the continuous phase of the emulsion in an amount of from 0.01 to 80% w/w based on the non-ionic surfactant, but which is such that the theoretical conductivity of the emulsion is less than the bulk conductivity of the emulsion; and
 - (e) water.

In a third aspect the present invention provides the use of a non-ionic surfactant and at least one polar or ionic or aromatic or conjugated compound in an amount of from 0.01 to 80% w/w based on the non-ionic surfactant to enhance the electrostatic charge imparted to droplets of a composition in the form of a water-in-oil or an oil-in-water emulsion on discharge from an aerosol spray device, which composition includes:

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- (a) at least one propellant in an amount of from 2 to 80% w/w;
- (b) optionally one or more solvents within the oil phase in an amount of up to 40% w/w; and

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and the amount of the polar or ionic or aromatic or conjugated compound being such that the theoretical. conductivity of the emulsion is less than the bulk conductivity of the emulsion. Program which had no

In a fourth aspect the present invention provides an aerosol spray which contains an electrically neutral composition in the form of a water-in-oil emulsion, an oil-in-water emulsion or a single phase ... composition, in which droplets of the composition on discharge from the aerosol spray device are imparted with a unipolar electrostatic charge, wherein the formulation of the composition and the material of the portion of the aerosol spray device with which the liquid comes into contact on spraying are selected such that 1.42 - 42 pol

- the difference between the Lewis base. i) component of the liquid and the Lewis base component of the material with which the liquid comes into contact on spraying is at least ± 5mJm⁻²; \$ 1.5
- the difference between the Lewis acid ii) component of the liquid and the Lewis acid component of the material with which the liquid comes into contact on spraying is at least ± 0.5 mJm⁻².5

The liquid droplets preferably have a charge to. mass ratio of at least $+/-1 \times 10^{-4}$ C/kg, more preferably at least $+/-2 \times 10^{-4}$ C/kg. The higher the 30 charge to mass ratio of the liquid droplets, the more effective the liquid droplets will be for their intended use, such as precipitating airborne particles and targeting insects. This charge level is considerably higher than the charge level which is 35

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achieved when spraying conventional liquid formulations from conventional aerosol spray device where charge devels of the order of +/- 1 x 10-5 to +/- 1/x/10 C/kg are obtained.

In the formulations of the present invention it is the combination of components (b) and (d) of the emulsion which improves the electron transfer through the emulsion with the charge being transferred from droplet to droplet through the emulsion at the interface between the disperse phase and the continuous phases and a second second

The various components (a) to (e) of the compositions of the present invention are discussed in turn below: 14 10 11 End to the ground of the control of the contr

Propellants

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One or more propellants are used in the composition of the invention in a total amount of from 2 to 80% w/w. Amongst the propellants that may be used are hydrocarbons and compressed gas, of which hydrocarbons are preferred.

Hydrocarbon propellants which may be used are acetylene, methane, ethane, ethylene, propane, nbutane, n-butene, isobutane, isobutene, pentane, pentene, isopentane and isopentene. Mixtures of these propellants may also be used . Commercially available propellants typically contain a number of hydrocarbon gases. For example, an odorised commercial butane, contains predominantly n-butane and some iso-butane. together with small amounts of propane, propene, pentane and butene.

Preferred hydrocarbon propellants include propane, n-butane, isobutane, pentane and isopentane, whilst the most preferred are propane, iso-butane and n-butane.

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Particularly preferred hydrocarbon propellants are mixtures of propane, n-butane and iso-butane.

whilst broadly the concentration of hydrocarbon propellant will be from 2 to 80% w/w, generally the concentration will be from 10 to 60% w/w, preferably 25 to 60% w/w and most preferably about 40% w/w.

when compressed gases are used as a propellant these will generally be carbon dioxide, nitrogen or air. Usually, they will be used at a concentration of 2 to 20% w/w, preferably about 5% w/w.

Non-ionic Surfactants.

Non-ionic surfactants for use in the present invention include mono, di and tri sorbitan esters, polyoxyethylene mono, di and tri sorbitan esters; mono and polyglyceryl esters; alkoxylated alcohols; alkoxylated amines; alkoxylated acids; amine oxides; ethoxylated/proproxylated block copolymers; alkoxylated alkanolamides; and alkoxylated alkyl phenols.

Particularly preferred are those surfactants which contain at least one alkyl, allyl or substituted phenyl group containing at least one C_6 to C_{22} carbon chain. Examples are esters with C_{10} - C_{22} fatty acids, preferably C_{12} - C_{18} fatty acids, particularly polyglycerol oleate and ethoxylated fatty alcohols, such as oleyl alcohol ethoxylated with two moles of ethylene oxide.

In some instances, the non-ionic surfactant may itself be combined with component (d). For example, when the non-ionic surfactant is polyglycerol oleate, the surfactant may contain small quantities of sodium or potassium oleate, for example is an amount of from 0.01 to 1% by weight. Greater quantities of such ionic compounds are generally not desirable and may

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result in the compositions not fulfilling the conductivity requirements of the compositions of the invention.

The concentration of the non-ionic surfactant is from 0.01 to 10% w/w, preferably 0.01 to 1% w/w.

Polar, ionic, aromatic or conjugated compounds

The polar or ionic or aromatic or conjugated compound which is included as component (d) in the compositions of the present invention may be selected from:

- a) alkali metal salts, alkaline earth metal salts, ammonium salts, amine salts or amino alcohol salts of one or more of the following compounds: alkyl sulphates, alkyl ether sulphates, alkylamidoether sulphates, alkylarylpolyether sulphates, monoglyceride sulphates, polyglyceride sulphates, alkyl sulphonates, alkylamine sulphonates, alkylaryl sulphonates, olefin sulphonates, paraffin sulphonates, alkyl sulphosuccinates, alkylamide sulphosuccinates, alkylamide sulphosuccinates, alkyl sulphocinnamates, alkyl sulphoacetates, alkyl phosphates, alkylether phosphates, acyl sarcosinates, acyl isothionates and N-acyl taurates;
- b) alkyl amidopropylbetaines, alkylamidobetaines, alkylamidosulphobetaines, alkylbetaines, aminimides, quaternary ammonium compounds and quaternary phosphonium compounds;
 - c) carboxylic acids, carboxylic acid salts, esters, ketones, aldehydes or amides of carboxylic acids containing from 6 to 30 carbon atoms;
 - d) diethyl orthophthalate, methylphenylcarbinyl acetate, α-methyl ionone, 4-hydroxy 3-methoxy-benzaldehyde, phenylethyl alcohol, dipropylene glycol, styryl acetate, n-butyl benzoate, isopropyl 4-

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hydroxybenzoate, isobutyl acetophenone, isopropyl acetophenone, nicotinic acid, benzoic acid, 2-naphthol, neopentyl benzene, naphthalene, toluene and fullerene.

Particularly preferred compounds in group (b) are alkyldimethylbenzyl ammonium chloride, octyltimethyl ammonium bromide, cetyltrimethylammonium bromide and dodecyltrimethylphosphonium bromide.

Particularly preferred compounds in group(c) are lauric, oleic, palmitic, ricinoleic and stearic acids, or the salts amides, esters, ketones or aldehydes thereof.

The concentration of component (d) is from 0.01 to 80% w/w, preferably from 0.01 to 30%, more preferably from 0.1 to 10% w/w based on the non-ionic surfactant, component (b). The amount of component (d) is selected so that the bulk conductivity is greater than the theoretical conductivity. In some cases, too great an amount of component (d) can result in the composition not fulfilling the conductivity requirement of the compositions of the invention.

Solvents

One or more solvents may be incorporated in the compositions of the invention in an amount of up to 40% w/w. Generally, the solvent will be water immiscible.

A wide range of solvent materials may be used, although care should be exercised to ensure that the solvent does not adversely interact with any active components of the compositions of the invention, such as insecticides.

Examples of solvents that may be used in the compositions of the invention include:-

liquid n-paraffins, liquid isoparaffins,

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cycloalkanes, naphthene-containing solvents, white spirit, kerosene, ester solvents, silicone solvents or oils, fatty acids, dialkyl phthalates, C₅-C₁₁ alcohols and fatty alcohols. Specific examples of these are as follows:

liquid n-paraffins - Norpar 12, Norpar 13 and Norpar 15 (available from Exxon)

liquid isoparaffins - Isopar G, Isopar H, Isopar L, Isopar M and Isopar V (available from Exxon).

Naphthene-containing solvents - Exxsol D40, Exxsol D60, Exxsol D80, Exxsol D100, Exxsol D110, Nappar 10, Solvesol 100, Solvesol 150, Solvesol 200 (available from Exxon)

Ester solvents - such as alkyl acetates, examples being Exxate 1000, Exxate 1300 (available from Exxon), and Coasol (available from Chemoxy International); c. Silicone solvent oils - Dow Corning 244, 245, 344

and 345 fluids,

Fatty alcohols - octanol, dodecanol, lauryl

alcohol, myristyl alcohol, cetyl alcohol, stearyl alcohol, cetostearyl alcohol, oleyl alcohol.

Preferred solvents are liquid hydrocarbon solvents, n-paraffins, and iso-paraffins.

Although the solvent is preferably incorporated as a level of from 1 to 20% w/w, more preferably the concentration will be in the range of from 2 to 10% w/w, most preferably about 5% w/w.

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Insecticidal Compositions

In one preferred agent of the present invention the compositions are insecticidal compositions which contain from 0.001 to 5% w/w of an insecticidal compound. A wide range of active ingredients may be used of which pyrethroids, particularly synthetic pyrethroids, chlorpyrifos, propoxur and diazinon are

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When synthetic pyrethroids such as lambda. cyhalothrin and bioresmethrin are used, generally they will be incorporated in concentrations of about 0.02% w/w or above. Whence the sectionado terrantos a terrological

Other synthetic pyrethroids such as cypermethrin, tetramethrin, permethrin and bioallethrin, will usually be incorporated to give a concentration of about 0.2%-0.5% w/w, or above.

Chloropyrifos, propoxur and diazinion will generally be incorporated to give concentration in the range of 0.5-0.9% w/w.s. in the series

... Preferably, insecticidal compositions of the invention will include an insecticide which functions primarily to knock down an insect, together with a second insecticide which functions primarily as a kill agent. An example of such a combination is the use of permethrin as a kill agent in a combination with tetramethrin as a knock down agent.

Optional Ingredients

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Various optional ingredients may be incorporated into the compositions of the present invention. For example, in order to maximise the effectiveness of the insecticidal activity of the compositions of the invention, synergists such as N-octylbicycloheptene dicarboximide and piperonyl butoxide may be included. at a concentration of from 0.5 to 1.5% w/w, most preferably about 1.0% w/w, for use in conjunction with pyrethroid insecticides.

In addition, other ingredients including corrosion inhibitors, such as 1-hydroxyethyl-2heptadecenyl imidazoline and/or sodium benzoate, preferably in a concentration from 0.01 to 0.5% w/w, preservatives and antioxidants, such as butylated

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hydroxytoluene, may be used as required. One or more fragrance components may also be included, according to the particular consumer requirements. It will be understood that certain fragrance components are components which may comprise component (d) of the compositions of the present invention and in this instance, such a component is not an optional ingredient.

Lewis Acid and Lewis Base Characteristics

In relation to the fourth embodiment of the invention an aerosol spray device and the electrically neutral composition contained therein have certain lewis acid and Lewis base characteristics which assist in imparting a unipolar charge to the liquid.

when two substances are brought together and then separated, an electrical charge is transferred from one to the other. This can occur for solid-solid separations, for solid-liquid separations and for liquid-liquid separations. When one of the components becomes airborne, the electrical charge can remain on the substance for a significant length of time as there is no place to ground the charge. The ability of substances to transfer their charge can be related to the characteristic Lewis acid (y') and Lewis base (y') values for the substance. All substances have characteristic values and these can be calculated indirectly from their component surface energies.

These surface energies when combined with a London-Van der Waal component (γ^{LN}) form what is known as the surface tension. This is easiest to measure at the interface between a drop of liquid on a solid substrate.

This equation used for these calculations comes from the approach used by Good and vanOss:

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 $\begin{array}{lll} (1 + \cos \theta) \gamma_L = 2 [\sqrt{(\gamma_a^{LW} \gamma_L^{LW})} + \sqrt{(\gamma_s^* \gamma_L^-)} + \sqrt{(\gamma_s^* \gamma_L^-)}] \\ & \text{Where } \theta \text{ is the contact angle that a drop of liquid makes with the surface.} \\ \end{aligned}$

 γ_e is the energy components for the solid. γ_L is the energy component for the liquid γ^{LM} Represents the London-Van der Waal's component.

 γ^{+} is the Lewis acid component (electron acceptor).

γ is the Lewis base component (electron donor).

Using three or more test liquids whose

characteristics are known, it is possible to solve this equation for the three unknowns, γ_s^{LW} , γ_s^{+} and γ_s^{-} . By using test solids whose characteristics are known, it is also possible to solve this equation to find the three unknown liquid surface energies. Thus it is

three unknown liquid surface energies. Thus it is possible to characterise a series of solids and liquids to form a series of Y and Y values. In the case of solids this series matches that of the

triboelectric series. Below is a non-exclusive list of solid substances that form part of this triboelectric series. The values are scaled such that water has a Y and Y of 25 mJm⁻². The top of the

list tends to become positive, when separated from a substance at the bottom of the list, which would

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become negative.

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Surface ಂದ್ಯ ಕ್ರಾಮಾರ್ ಚಿಕ್ಕಾರ್ ಎಂದು ಕಟ್ಟಿಸಿದ್ದಾರೆ. ಬಿಡಿಕಾರ್ ಪ + Glass Acetal 900P NC-10 Milesan - 15 has buttered at the control of the celluloid - let ame separation and at the PMMA TO DE LEAS TO THE FRONT COLORS OF 12 TO SHEET OF THE SHEET OF Nylon Treversel 189 1898 pl fillselters to 8.4 PVC Polyester 2002-2 5 5 5 0.1 Polyethylene Cartinate the contract of the second of the Later Brown Car Williams East Bur

Liquids also have a value of Y and Y that can be measured. When liquids contact a solid of known Y and γ^* on aerosol spraying, and the γ^- of the liquid is greater than, or less than that of the solid by 520 - 40 mJm⁻², most preferably 15 mJm⁻², or a ytogreater than, and the or less than that of the solid by 0.5 mJm preferably. 1 mJm⁻², most preferably 2 mJm⁻², the liquid has a tendency to charge to '± 1 x 10 C/kg. A common material from which inserts for aerosol actuators are made is Acetal 900P NC-10, which has a γ of 15 mJm⁻².

Aerosol Spray Devices

The compositions of the present invention, when .. sprayed through conventional aerosol spray heads; form into droplets which are imparted with a unipolar charge of at least $+/-1 \times 10^{-4}$ C/kg.

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It is possible to impart even higher charges to the liquid droplets by choosing aspects of the aerosol device including the material, shape and dimensions of the actuator, the actuator insert, the valve and the dip tube and the characteristics of the liquid which is to be sprayed, so that the required level of charge is generated as the liquid is dispersed as droplets.

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A number of characteristics of the aerosol system increase double layer charging and charge exchange between the liquid formulation and the surface of the aerosol system. Such increases are brought about by factors which may increase the turbulence of the flow through the system, and increase the frequency and velocity of contact between the liquid and the internal surface of the container and valve and actuator system.

By way of example, characteristics of the actuator can be optimised to increase the charge levels on the liquid sprayed from the container. A smaller orifice in the actuator insert, of a size of 0.45mm or less, increases the charge levels of the liquid sprayed through the actuator. The choice of material for the actuator can also increase the charge levels on the liquid sprayed from the device with materials such as nylon, polyester, acetal, PVC and polypropylene tending to increase the charge levels. The geometry of the orifice in the insert can be optimised to increase the charge levels on the liquid as it is sprayed through the actuator. Inserts which promote the mechanical break-up of the liquid give better charging.

The actuator insert of the spray device may be formed from a conducting, insulating, semi-conducting or static-dissipative material.

The characteristics of the dip tube can be optimised to increase charge levels in the liquid sprayed from the container. A narrow dip tube, of for example about 1.27mm internal diameter, increases the charge levels on the liquid, and the dip tube material can also be changed to increase charge.

Valve characteristics can be selected which increase the charge to mass ratio of the liquid

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product as it is sprayed from the container. A small ... tailpiece orifice in the housing, of about 0:65mm; increases the chargento mass ratio during spraying. A reduced number of tholesain the stem, for example 2 x: 0.50mm; lalso increases the charge during spray. The presence of a vapour phase tap helps to maximise the charge levels, a larger orifice vapour phase tap of, for example, about 0.50mm to:1.0mm generally giving higher charge levels.

The liquid droplets sprayed from the aerosol spray device will generally have diameters in the range of from 5 to 100 micrometres; with a peak of : " droplets of about 40 micrometres.

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Preparation

The compositions of the present invention may be prepared by standard techniques which are well known in the art. For example, components (b) to (d) may be mixed together to form the solvent phase. This solvent phase is then mixed with water to produce an emulsified concentrate which is then filled into cans and blended with the propellant. Alternatively, the concentrate and the propellant may be filled into the and the second s cans simultaneously.

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Conductivity the paint and product and specifical

The theoretical conductivity of an emulsion, o, can be calculated from measurements of the actual conductivity of the external phase and the internal phase, according to the following equation:

$$\sigma_{\rm c} = \sigma_{\rm c} (1-3\phi (\sigma_{\rm p} \sim \sigma_{\rm c})/(\sigma_{\rm p} \sim 2\sigma_{\rm c}))$$

o = theoretical conductivity of the where emulsion and the transfer of the second

 $\sigma_c = measured conductivity of the$ separated external phase

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opt≡the separated internal phase

The bulk conductivity can be determined by ; experimentation: Component (d) of the compositions of the present invention serves to enhance the actual conductivity of the emulsion and thus the bulk conductivity of the emulsion is higher than the theoretical conductivity calculated according to the above formula. Preferably the difference between the theoretical conductivity of the emulsion of the emulsion is at least $+0.5\mu S_{\odot} cm_{i}$ preferably at least $74\mu\mathrm{S}$ cm, more preferably at least + $6\mu\mathrm{S}$ cm. Component (d) in the compositions of the present invention thus improves the electron transfer through the emulsion with the charge being transferred from droplet to . droplet through the emulsion at the interface between the disperse phase and the continuous phase.

The present invention will be further described with reference to the following mon-limiting Examples.

At words of the treatment of

In the control of the

Method for predicting the theoretical conductivity of an emulsion through the measurement of the conductivity of the individual phases:

- 1. Calibrate the conductivity cell by measuring a solution of known conductivity. The conductivity cell comprises a pair of platinum electrodes, held apart and attached to the inside of a glass tube of approximately 1cm internal diameter.
 - Using the same cell, measure the conductivity of the bulk emulsion according to the invention, whilst ensuring that the

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emulsion isostatic and homogeneous prior to taking the measurement. 333 Determine whether the continuous phase of the emulsion is water or oil. 4.0 Separate the two phases of the emulsion by Teither gravimetric or centrifugal (). () Separation: Isolate the phases and measure the conductivity of each phase in the Nocalibrated cell, and waster that the first of the control of the 5. 5. Use the equation given below to determine -the theoretical conductivity of the bulk do and emulsion. The second of the second second second second 6. The difference between the theoretical conductivity and that obtained directly from the measurements is the contribution to the bulk conductivity=due to the *emulsion** with the temple of the temple o on or droplets of law when the prices agreed was labor $\omega = \sigma_c^2 = \sigma_c^2 (1-3\phi(\sigma_c + v\sigma_c)/(\sigma_c + 2\sigma_c)) = 2\sigma_c + \sigma_c +$ where of theoretical conductivity of the Table to the company of emulsion and the control of the control $\sigma_{\rm c} = \text{measured conductivity of the}$ ' separated external phase σ_p = measured conductivity of the separated internal phase and a set

· φ = volume fraction of the external

na phase and a same of meaning as a sign of

Measurement of Electrostatic Charge

The charge to mass ratio of the compositions of Examples 4 to 50 was measured using a standard aerosol can with a valve insert made from polyoxymethylene according to the design shown in accompanying Figure 1 e de la tradición de la companyo 9 in which

Fig. la is a plan view;

Fig. 1b is a sectional view; and

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Fig. 1c is a inverted plan view of the valve insert.

EXAMPLE 1 (Comparative) - Problems 100

5	.!	a backs intact
	An emulsion was prepared from the	
	ingredients: //	er de ferriera
		. <u>*</u>
	Ethoxylated (7EO)alcohol(C_{12} - C_{15})	
10	Deionised water	47 v/v
	Decane	55 v/v
	Actual (measured) conductivity	1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 - 1960 -
	of the bulk emulsion	
	σ_{c}	16.6 μ S cm
15	$\sigma_{_{\mathbf{P}}}$.	4.1 μ S cm
	Assuming of =	0.5
	SO THE LOSE	8.2 µS cm
	Difference (conductivity due the en	nulgion
	• • • • • • • • • • • • • • • • • • •	

droplets) = 7.4 - 8.2 = -0.8 \u03cm.

This composition did not contain any component
(d) and as can be seen the bulk conductivity of the emulsion is less than the theoretical value.

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An emulsion was prepared from the following is validate que to, and all all and ingredients:

Ingredient 5 Ethoxylated (7EO) alcohol (C12-C15) 0.24 w/v 3% w/w of the Sodium lauryl sulphate non-ionic (30% active) services a legal probability of surfactant 18. 29. 47. v/v. 50 Deionised water 10 53 v/v ** Decane Actual (measured) conductivity of the bulk emulsion $22.3~\mu\text{S}$ cm $39.4~\mu\text{S}$ cm σ_c 4.0 µS cm 15

 $\sigma_{\mathbf{p}}$ 4 80 p5.5% to lead Assuming φ = 14.1 μ S cm o L

Difference (conductivity due the emulsion: droplets) = $22.3 - 14.1 = 8.2 \mu \text{S} \text{ cm}$. Clarity of the Alberta Company of the Company the contract that will be the stress of the first of the contract of the contr

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EXAMPLE 3 (Comparative)

				•	
	• :. • —		41	·. ··	
- An emulsion	.was pr	epare	1 from	the	following
ingredients:om	1 2 3 S	* . "	ాంద్ జ ల్ -	: i - :	———— ఇంజారణ ఎతంతో ఈ ఎగుగుమ్

5	Ingredient	%			
	Ethoxylated (7EO)alcohol($C_{12}-C_{15}$)				
	Sodium lauryl sulphate:	6% w/w of the			
		non-ionic			
		surfactant			
10	Deionised water	47 v/v .:			
	Decane	53 v/v			
	Actual (measured) conductivity				
	of the bulk emulsion of the bulk emulsion	23''3' //S' cm			
	σ _e ,	83.0 %S cm			
15	G G	9.0 μS cm			
	Accumina	0.5			
		30.3 μS cm			
	Difference (conductivity due the emu	lsion:			
	droplets) = 23.3 - 30.3 = -7.0 μ S cm.				
20	This composition contains a large amount of				
	component (d) and as can be seen the bull	k conductivity			
	of the emulsion is less than the theoret:	ical value			

EXAMPLE 4

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	Ingredient	<u>4</u>
	Ethoxylated (SEO) alcohol	_ <u></u>
	(C ₁₂₋₁₅) incorporating benzalkonium	
_0	chloride (50% active) at 2% w/w in	
30	surfactant	0.24% w/v
·	Deionised water	47% v/v
	Butane 40	53% v/v

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This formulation when made up as an aerosol and sprayed though the physical valve/actuator combination

The second second

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described above produced a mono-polar charge on the sprayed droplets of +1.65 x 10 - C/Kg.

The same formulation was prepared substituting decane for butane in order that the conductivity could be measured.

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Actual (mea	asured) conductivity	y	ur.A.	
of the bull	k emulsion	\$ 73°C	15.7 μ S	cm
$\sigma_{f c}$		•	40.2μS	cm
σ _p		\$2.77 TH \$ 1.77	3.3µS	zm
Assuming 0	=		0.5	
σ	er der andtrick in 18	استواد المستواد	13.7 μ \$	cm
Difference	(conductivity due	to the 🗀 🦥	77.1 4.1	
emulsion di	ronlets) = 15.7 - 13	.7= 2.04S	cm ·	

EXAMPLE 5

Marie Contraction

	Formulational and sufferent sign from	
	Ingredient:	% w/w
20	Solvent	
	Fragrance component	
	Butylated hydroxytoluene	0.7013
	Polyglyceryl oleate	0.30
	Deionised Water	58.99
25	Butane 40	35

The solvent used and the fragrance component that can be used in the above formulation are illustrated in the following examples:

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Charge/Mass (x10-4 C/kg)

Fragrance Component

Solvent

Example

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-2.2

diethyl-o-phthalate

Isopar E Isopar E Isopar G Isopar G

styrallyl acetate α-methyl ionone

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rom I Ltd.	
PT 0	

The Isopar E, G and L range of solvents can be obtained

-2.2 2.5

-2.4

phenylethyl alcohol

Isopar E Isopar L

dipropylene glycol

-1.7

Litsea Cybeba

heptane pentane

Lilial

vanillin

-2.3

6.1-7.6

The fragrance components used were obtained from Robertet

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EXAMPLE 13

Form	<u>ılat</u>	<u>ion</u>	<u>2</u>
------	-------------	------------	----------

	Ingredient	% w/w
5	Solvent	5.0
	Aromatic component	0.70
	Butylated hydroxytoluene	0.013
	Pòlyglyceryl öleate	0.30
•	Deionised Water	58.99
10	Butane 40	35

A CHANGE CAS

The solvent used and the aromatic component can be used in the above formulation are illustrated in the following examples:

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EXIM

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'Aa			•	-	•	-					THE SHIELDS LETS
96	, _		_					_			An interpretation
Charge/Мавв (x10 → C/kg)	3.0	-2.5	-3.0	+1.6	+1.7	+1.2	 	÷1.9	ः <u>।</u> फ़	5.4	i G Malov grani i mani go t
			a						,		SEARCH OF BUILDINGS OF LINE WELL
	17		isopropyl-4-hydroxybenzoate	<u>a</u>	ne	I	365			* *	
护		. 3	×X	, E	eno	A c	5 11		i Ligh	^Am	rioti a il composito di Cara
19C	nce	t e	dro	phe	oph				ene		20
oğu,	gra	zoa	-hy	eto	cet	Ö	• .		enz	:- •	/09
දි	fra	ben	Ţ-7	åC	.⊣ a	aci	0]		<u>н</u>	ene	o o
tte	ŏ	71	, go	[Y]	Vido	Ċ.	hth	Je.	nty	hal	n Go rn de Company de Christian de Christia
Aromatic Component	Cleanox fragrance	n-butyl benzoate	isopr	isobutyl acetophenone	isopropyl acetophenone	benzoic acid	2-naphthol	toluene	neopentyl benzene	naphthalene	fullerene C60/
	150										
		臼	J	[22]	41	۵.	>	. ტ		ບ	ප
/ent	/eBC	ar	ar	ar	ane	ane	ar	ar	ane	ar	н
Solvent	Solvesol	Isopar	Isopar	Isopar	heptane	pentane	Isopar V	Isopar G	Pentane	Isopar C	Isopar G
	<u></u>	F-7		, -		14	Н	Н	щ	H	
Example	13	14	15	16	17	18	19	20	21	22	23

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EXAMPLE 24

Formulation 3

<u>&</u> Ingredient Ethoxylated (5EO) alcohol (C_{12-15}) incorporating sodium laureth sulphate (30% active) 0.24% W/V at 4% w/w in surfactant 47% V/V Deignised water 53% V/V Butane 40

This formulation, when made up as an aerosol and sprayed through the physical valve/actuator combination described above produced a mono-polar charge on the sprayed droplets of -1.1 x 10 d C/kg.

The same formulation was prepared substituting decane for butane. The formulation had a voof 26.2 ${\mathfrak m}{\mathbb J}{\mathfrak m}^{-2}$.

An acetal 900P NC-10 insert in the spray head had a γ^{-} of 15 mJm⁻².

The difference between these Lewis acid values $= 26.2 - 15 = 11.2 \text{ mJm}^{-2}$

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Ingredient	M/M8	&w/w	M/M&	
	Example 25	Example 26	Example 27	
Bioallethrin	0.194			82 өтбілеуу
Bioresmethrin	900		7 · 1 / 4	0.194
ВНТ	970.0	0.036	0.036	0,036
חסויין האפועוריין ויין	0.02	0.02	0.02	0.02
Olein and	006.0	006.0	0.900	0.900
Derfine	60°0	0.180	0.045	0.023
Normar 13	0.100	0.100	0.100	00.100
Delonied water	7,500	7.500	7.500	7,500
HSS	51.16	51.07	51.205	51.227
Charge/mage/vio-4	40.000	40,000	40.000	40.000
(6y/) 0TV) ggpii />6	-0.75	-0.72	-0.63	06.0-
Ingredient	8w/w	M/MB	m/mg	8w/w
	Example 29	Example 30	Example 31	Example 25
Teric 12A2	0,800	1.000		
Oleic acid	0.400	0.400	0 0	0.700
Norpar 13	7 500		0.400	0.400
Deionised water	000	7.500	7.500.	7.500
H55	31.300 40.000	51.100	51.300	51.400
Charge/mass(x10-4 C/kg)	-1.02	40.000	40.000	40.000
	Ţ	000	-0.816	-0.816

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Ingredient	8w/w	8W/W	#/wg	;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;
	Екапрје 33	Example 34	Example 35	
Teric 12A2	0,600	0.500	1.000	0.000
	0.400	0.400	0.200	
	7.500	7.500	7,500	201
water	51.500	51.600	51,300	ý.
	40.000	40.000	40.000	Mary Control of the C
Charge/mавв(х10-4 С/kg)	-1.596 -1.596	-0.966	-1.53	#1. **
	8w/w	8w/w	, si	
Topa	катріе 36	Example 37	930 34E	10 % 10 % 10 % 10 %
	1.000	1,000	in in	-
Oleic acid	0.400	008.0	:. 	
	0,100	0.100	, <u></u>	90°.4
	7,500	7.500		
water	51,100	50.700	· ·,	*
-	40.000	40.000		·
' c/kg)	-0.57	-0.738	,	
		18,0 3-11 3	٠.	·.
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Ingredient	8w/w	%/m%	8w/w
	Example 38	Example 39	Example 40
Teric 12A2	1,000	1.000	1.000
Lauric acid	0,100	0.200	0.400
Norpar 13	7,4500	7.500	7.500
Deionised water.	51.400	51.300	51,100
H55	40.000	40.000	40.000
Charge/mass(x10-4 C/kg)	-0.532	-0.578	-0.574
Ingredient	8w/ws 61	8W/W	
	Example 41	Example 42	-
Teric 12A2	1.000	1.000	
Palmitic acid	0.10.	0020	:
Norpar 13	7,500	7,500	
Deionised water	51.400	51,300	
H55.	40.000	4,0 .000	
Charge/mass(x10 C/kg)	-0.502	-0.704	
		· . , ;	

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Ingredient	M/M%	8m/m8	M/M%	*/**
	Example 43	Example 44	Example 45	Example 46
Teric 12A2	0.700	0.720	0.700	0.700
Oleic acid	0.500	0,400	0.400	0.400
Nordar 13	7.500	7,500,	7.500	7.500
water	4,1,300	31.38	51,400	51.300
.H55	F.,	: :	t	40.000
.H46.	50.000	000-09	40.000	ı
Charge/mass(x104 C/kg)	(J-1), 3,9	24.2.2)	-0.71	-1.65
	960 -	· · · · · · · · · · · · · · · · · · ·		
Ingredient	8w/w	To a Tallings		
	Example 47	W. 200		
Teric 17A2	0.85			
Oleic acid	0.35	÷ā		
	5.00-	764.88	N.4. 6.8	
Deionised water	33,80	en. en en en	 	
, H46:	000.09	20 12 13	·· .	
Charge/mass(x10-' C/kg)	7-4.8	9		
		1.7 1.73 1.7		
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	£.	•	

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Ingredient	8W/W	W/W8
	Example 48	Exampl
Norpar 13	2,00	5.00
Bioallethrin	0.25	0.25
Bioresmethrin	0.05	0.05
BHT	0.02	0.02
Deionised water	33.28	33,58
Teric 17A3	0.60	0.35
Crill 45	0.40	0.35
Pine, Fragrance	0.10	0.10
Oleic acid	0.30	0.30
H46	60.00	60.00
Charge/mass(x10-4 C/kq)	-1.41	-1 34

Ingredient		A/MS
		Example
Bioallethrin		0.209
Bioresmethrin		0.039
BHT		0.005
Polyglycerol oleate		
(containing from 0.01		
to 1% by weight of		
sodium or potassium oleate)	;, 1	0.900
Perfume	yes N N N M N M N N	0.100
Norpar 13	157	7.500
Deioniged water	·	51.247
н55	- 1	40.000
Charge/mass(x10-4 C/kg)	71 12	-1,59
-		

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Comparative	Example' A	-	-		. :	4.				. , .			
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3d I		÷i r)] e	B		31 <u>y</u>	лие	3.r	ij		4	Je/	
Ingredient		Teric 12A2	Bioallethrin	Bioresmethri	Ē	Polyglycerol	Perfume	Norpar 13	Delonised wa	ις	Crill 45	Charge/mass	
H		Te	Bi	Bi	BHT	М	Pe	NC	Ď	H55	ŭ	ដ	

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Ingredients and Availability

	Oleic acid¹ :		Technical (Croda Chemicals)
	Lauric acid¹ :		Reagent Grade (BDH)
5	Palmitic acid¹ :		Reagent Grade (BDH)
	Teric 17A3 ² :		C ₁₇ alcohol with 3 moles of
			ethylene oxide (Orica)
	Teric 17A2 ² :		C ₁₇ alcohol with 2 moles of
			ethylene oxide (Orica)
10	Teric 12A2 ² :		C ₁₂ alcohol with 2 moles of
	э		ethylene oxide (Orica)
	Crill 45 ²		Sorbitan trioleate (Orica)
	Polyglycerol oleate ² :		Croda Chemicals
	BHT ⁴ :		Butylated hydroxytoluene
15	÷		(Orica)
	Norpar 13 ⁵ :		liquid n-paraffin (Exxon)
	Bioallethrin ³ :		93% w/w (Agrevo)
	Bioresmethrin ³ :	<u> </u>	93% w/w (Agrevo)
	H46 ⁶ :		16% w/w propane/butane blend
20			(Boral)
	H55 ⁶ :		26% w/w propane/butane blend
	• •		(Boral)
	1: Ionic Compound (d)	,· :	2: Non-ionic surfactant
	3: Insecticide	•	4: Antioxidant
25	5: Solvent		6: Propellant

STRAFT SECTION

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no CLAIMS: General for gold party and Mild of

- An electrically neutral composition in the form of a water-in-oil-or an oil-in-water emulsion, in which droplets of the emulsion on discharge from an aerosol spray device are imparted with a unipolar electrostatic charge, which composition comprises:
 - (a) at least one propellant in an amount of from 2 to 80% w/w; plant the result of the action of the second of the
 - (b) at least one non-ionic surfactant in an amount of from 0.01 to 10% w/w;
 - (c) optionally one or more solvents within the oil phase in an amount of up to 40% w/v;
- (d) at least one polar or ionic or aromatic or linearly conjugated compound which is attracted to the interface between the disperse phase and the continuous phase of the emulsion in an amount of from 0.1 to 80% w/w based on the non-ionic surfactant, but which is such that the theoretical conductivity of the emulsion is less than the bulk conductivity of the emulsion; and
 - (e) water.
- 2. A composition as claimed in claim 1 wherein the difference between the theoretical conductivity of the emulsion and the bulk conductivity of the emulsion is at least + 0.5μS cm.
- 3. A composition as claimed in claim 2 wherein the difference between the theoretical conductivity of the emulsion and the bulk conductivity of the emulsion is at least + 4μ S cm.
- 4. A composition as claimed in claim 2 wherein35 the difference between the theoretical conductivity of

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the emulsion and the bulk conductivity of the emulsion is at least + 6μ S cm.

A Composition as claimed in any one of the preceding claims wherein at least 90% by volume of the droplets of the disperse phase within the emulsion have an average diameter of less than 60 mm. ការស៊ុន ស្រែស្រាស់ មាន ស្រែស្រាស់ ស្រែស្រាស់ ស្រែស្រាស់ ស្រែសាស្រាស់

A composition as claimed in claim 5 wherein at least 90% by volume of the droplets of the disperse phase within the emulsion have an average diameter in the range of from 20 to 40 mes a line is a line

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- 7. A composition as claimed in any one of the preceding claims wherein at least one non-ionic 15 surfactant is selected from mono, di and tri sorbitan 🕦 esters, polyoxyethylene mono, di and tri sorbitan esters; mono-and polyglyceryl esters; alkoxylated alcohols; alkoxylated amines; alkoxylated acids; amine oxides; ethoxylated/proproxylated block copolymers; 20 alkoxylated alkanolamides; and alkoxylated alkyl phenols.
- 8. A composition as claimed in claim 8 wherein the ionic surfactant contains at least one alkyl, 25 allyl or substituted phenyl group containing at least THE CANCEL SERVICE STREET one C₆ to C₂₂ carbon chain.
- A composition as claimed in any one of the preceding claims wherein component (d) is selected '-30 from
 - alkali metal salts, alkaline earth metal a) salts, ammonium salts, amine salts or amino alcohol salts of one or more of the following compounds: alkyl sulphates, alkyl ether sulphates, alkylamidoether

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sulphates, alkylarylpolyether sulphates, monoglyceride sulphates, polyglyceride sulphates, alkyl sulphonates, alkylamine sulphonates, alkylaryl sulphonates, olefin sulphonates, paraffin sulphonates, alkyl sulphonates succinates, alkylether sulphosuccinates, alkylamide sulphosuccinates, alkylasulphocinnamates, alkyl sulphoacetates, alkylaphocinnamates, alkylaphocetates, alkylaphocetates,

- b) alkyl amidopropylbetaines, alkylamido-sebetaines, alkylamidosulphobetaines, alkylbetaines, aminimides, quaternary ammonium compounds and quaternary phosphonium compounds;
- c) carboxylic acids, carboxylic acid salts,
 esters, ketones, aldehydes or amides of carboxylic
 acids containing from 6 to 30 carbon atoms;

 d) acids thyl orthophthalate, methylphenylcarbinyl
 acetate, α-methyl ionone, 4-hydroxy 3-methoxybenzaldehyde, phenylethyl alcohol, dipropylene glycol,
 styryl acetate, n-butyl benzoate, isopropyl 4hydroxybenzoate, isobutyl acetophenone, isopropyl
 acetophenone, nicotinic acid, benzoic acid, 2-napthol,
 neopentyl benzene, naphthalene, toluene and fullerene.
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 10. A composition as claimed in any one of the preceding claims wherein component (d) is present in the composition in an amount of from 0.01 to 30% w/w, preferably 0.12 to 10% w/w based on the weight of component (b).
 - 11. A composition as claimed in any one of the preceding claims wherein the droplets formed on discharge from an aerosol spray device have a charge to mass ratio of at least +/- 1 x 10⁻⁴ C/kg, preferably at least +/- 2 x 10⁻⁴ C/kg.

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12 12 A Composition as claimed in any one of the preceding claims which is an insecticidal composition which includes one or more insecticides therein in an amount of from 0.001 to 5% w/w. There have been all the

FOR SIMPLE ENGRADORED TO SEE TO SEE TO SEE TO SEE THE SEE TO SEE THE S 13. A method of enhancing the unipolar charge which is imparted to droplets of an emulsion on discharge from an aerosol spray device in which the

droplets are formed from an oil-in-water or a waterin-oil emulsion composition which comprises:

(a) at least one propellant in an amount of from 2 to 80% w/w:ಏನ ಸಂಭಾವನ ಮಾನಾಯದ ಬಾಗುತ್ತಿದ್ದಾಯಕಾಲಕಾಗು ಇತ್ತಿದ್ದಾರ ಕ್ಷಮ

at least one mon-ionic surfactant in an and amount of from 0.01 to 10% w/w;

(c) optionally one or more solvents within the oil phase in an amount of up to 40% w/w; productions

(d) at least one polar or ionic or aromatic or conjugated compound which is attracted to the interface between the disperse phase and the continuous phase of the emulsion in an amount of from 0.01 to 80% w/w based on the nontionic surfactant, but which is such that the theoretical conductivity of the

emulsion is less than the bulk conductivity of the emulsion; and

only (e) water. The lower in the months of the con-ក្រុម ព្រះបាន ប្រជាជាធិប្បាយ នេះ ប្រាស់ នេះ ប្រាស់ មាន ប្រាស់ ប្រាស់ ប្រាស់ ប្រាស់ ប្រាស់ មាន ប្រាស់ មាន ប្រាស

14. The use of a non-ionic surfactant and at which least one polar or ionic or aromatic or conjugated or compound in an amount of from 0.01 to 80% w/w based on the non-ionic surfactant to enhance the electrostatic charge imparted to droplets of a composition in the form of a water-in-oil or an oil-in-water emulsion on discharge from an aerosol spray device, which composition includes:

(a) at least one propellant in an amount of from

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\$2 to 80% w/w; in a rest on garge lorence in

- (b) optionally one or more solvents within the oil phase in an amount of up to 40% w/w; and of the control of the
 - water;

and the amount of the polar or ionic or aromatic or conjugated compound being such that the theoretical conductivity of the emulsion is less than the bulk conductivity of the emulsion and the conductivity of the conductivity of the conductivity of the emulsion and the conductivity of the conductiv

- 10 15. An aerosol spray which contains an electrically neutral composition in the form of a water-in-oil emulsion, an oil-in-water emulsion or a single phase composition, in which droplets of the composition on discharge from the aerosol spray device are imparted with a unipolar electrostatic charge, 15 wherein the formulation of the composition and the material of the portion of the aerosol spray device with which the liquid comes into contact on spraying are selected such that
 - the difference between the Lewis base i) component of the liquid and the Lewis base component of the material with which the liquid comes into contact on spraying is at least $\pm 5 \text{mJm}^{-2}$; and/or:
 - the difference between the Lewis acid ii) component of the liquid and the Lewis acid component of the material with which the liquid comes into contact on spraying is at least \pm 0.5 mJm⁻².
 - 16. An aerosol spray device as claimed in claim 15 wherein the difference in i) is at least ± 10mJm⁻² and/or the difference in ii) is at least \pm lmJm⁻².

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17. An aerosol spray device as claimed in claim 15 or 16 wherein the difference i) is at least \pm 15mJm⁻² and/or the difference in ii) is at least \pm 2 mJm⁻².

18. An aerosol spray device as claimed in any one of claims 15 to 17 wherein the composition contained therein is a composition as claimed in any one of claims 1 to 14.

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ABSTRACT

COMPOSITIONS

An electrically neutral composition in the form of a water-in-oil or an oil-in-water emulsion, in which droplets of the emulsion are imparted with a unipolar electrostatic charge on discharge from an aerosol spray device.

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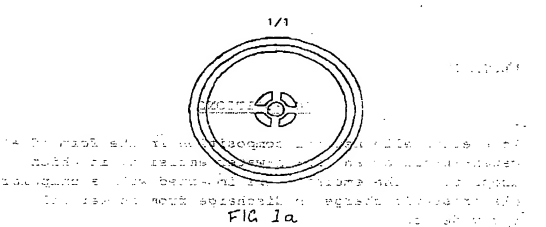




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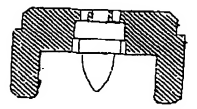
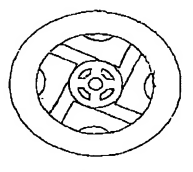


FIG 16



F: G 1c

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